Simulating multi-substrate diffusive transport in 3-D tissues with BioFVM

Samuel H. Friedman, Ahmadreza Ghaffarizadeh, and Paul Macklin, *USC PSOC Member*

Abstract—To simulate the spatiotemporal distribution of chemical compounds, we present BioFVM, an open-source reaction-diffusion equation solver using finite volume methods with motivation for biological applications. With various numerical solvers, we can simulate the interaction of dozens of compounds, including growth substrates, drugs, and signaling compounds in 3-D tissues, with cells by treating them as various source/sink terms. BioFVM has linear computational cost scalings and demonstrates first-order accuracy in time and second-order accuracy in space. Beyond simulating the transport of drugs and growth substrates in tissues, the ability to simulate dozens of compounds should make 3-D simulations of multicellular secretomics feasible.

I. INTRODUCTION TO THE TYPE OF PROBLEM IN CANCER

Tissues are filled with various chemical compounds, including signaling and other factors that regulate how cells move, grow, and die, depending on the concentration and/or gradient of any and all of these compounds. In order for cancer cells to survive and grow, they need to obtain oxygen and other nutrients released from blood vessels, and change phenotype based upon signaling factors released by other cells and the vasculature. These chemical substances move through tissues by diffusion, and are impacted by uptake by tumor and other cells and reaction terms (e.g., decay). (In some tissues, advection is also dominant; these effects are not considered in this version.) These transport processes can be used to model how chemotherapeutic drugs reach their intended targets: susceptible cancer cells. By performing computational simulations of the movement of dozens of various chemical substances and of cellular chemical uptake and secretion rates, we can test hypotheses that can control the overall growth of cancer cells and enable 3-D simulations of multicellular secretomics.

Simulating this motion requires solving a system of reaction-diffusion equations for a vector \( \vec{\rho} \) of substrates:

\[
\frac{\partial \vec{\rho}}{\partial t} = \vec{D} \odot \nabla^2 \vec{\rho} - \vec{\lambda} \odot \vec{\rho} + \vec{S} \odot (\vec{\rho}_T - \vec{\rho}) - \vec{U} \odot \vec{\rho} + \sum_{\text{cells } k} \delta(\vec{x} - \vec{x}_k) \left[ \vec{S}_k \odot (\vec{\rho}_{T,k} - \vec{\rho}) - \vec{U}_k \odot \vec{\rho} \right]
\]

where \( \vec{D} \) and \( \vec{\lambda} \) are the diffusion and decay coefficients, \( \vec{S} \) and \( \vec{U} \) are the bulk source and uptake terms, and \( \vec{S}_k \) and \( \vec{U}_k \) are cell-centered source and uptake terms. The \( \odot \) symbol denotes the Hadamard (element-wise) product between two vector terms, \( \delta \) is the Dirac delta function, and \( \vec{\rho}_T \) represents target densities of the substrates.

We present here BioFVM [1], an open-source software package written in C++ to efficiently solve this system of equations on multicore desktop computer and single supercomputer nodes. Using the finite volume method (FVM) [2], this software fills a niche and focuses on the biological transport processes of interest to cancer modelers. BioFVM is intended to work in concert with multicellular modeling packages (e.g., Chaste [3], CompuCell3D [4], PhysiCell [5]) to simulate how cells change their phenotypes as they interact with other cells and the biochemical microenvironment. Combining BioFVM with an agent-based code enables a user to simulate the growth, survival, and death of not only cancer cells, but of multicellular processes in general.

BioFVM is open source under the Apache 2.0 license. It can be downloaded for free at:


A series of tutorials is included with every download. The method is described in detail in [1].

II. ILLUSTRATIVE RESULTS OF APPLICATION OF METHODS

BioFVM obtains first-order accuracy in time, second-order accuracy in space, and linear scaling of cost in terms of each of the number of substrates, voxels, and time steps [1]. Simulating additional substrates does not significantly increase the computational cost of a simulation. Fig. 1 demonstrates the computational cost scaling when increasing

Figure 1: Number of substrates vs. execution time (seconds) for a simulation with otherwise identical conditions.
the number of substrates from 1 to 100.

Fig. 2 shows an example simulating oxygen and glucose transport in a liver lobule. We marked central veins in histological images of liver tissues, estimated the corresponding lobular geometry, generated sinusoids, added hepatocytes, and imported tissue into BioFVM with a 5 mm³ computational domain (5 mm × 5 mm × 200 μm). We simulated oxygen and glucose release by the portal triads, diffusion, and uptake by hepatocytes. The desktop simulation used 625,000 voxels (20 μm resolution) with 900,000 cells. See [1] for further examples and test results.

III. QUICK GUIDE TO THE METHODS

We discretize the spatial simulation domain into voxels to solve the reaction-diffusion equation with the finite volume method. BioFVM currently supports uniform Cartesian meshes, where the FVM reduces to a finite difference scheme; support for non-Cartesian Voronoi meshes is expected in a future release.

We several techniques to speed up numerical computations. We use operator splitting [6, 7] to solve the bulk source/sink terms first, the cellular source/sink terms second, and diffusion/decay terms third. The splitting is O(Δt) accurate and stable when the individual solvers are first-order accurate and stable [6]. Moreover, it simplifies development, allows efficient solvers to be tailored to each term, and holds the door open to new solvers (e.g., advection) without significant change to the overall code.

We solve the bulk and cell-centered source/sink terms using the (1st-order, stable) backwards Euler scheme. In each time step, the source/sink terms are independent across space. Thus, parallelize their solutions using OpenMP. See [1] for the full details of the discretization.

We divide the 3-D diffusion/decay terms into a series of 1-D equations, one for each dimension [6]. After this splitting, a strip of voxels (e.g. all voxels with the same yj and zk values, but different x values) is independent of any other strip; discretizing the diffusion and decay terms yields a tridiagonal linear system, which can be directly by using the Thomas algorithm [8]. Because each x strip is independent, we can distribute and run many instances of the Thomas solver across the processor cores with OpenMP, allowing us to easily parallelize the x diffusion problem. We solve along each of the y and z dimensions similarly.

To further optimize the code, we used some computational techniques that better utilize modern computer architecture, trading memory for computational speed wherever possible. Rather than solving the PDE for each chemical substrate sequentially (which requires 10× more computational time to solve for 10 substrates than solving for one substrate), we discretized the solvers to operate on entire vector of substrates simultaneously, thus taking advantage of SIMD and similar hardware optimizations on modern CPUs. We used BLAS operations (e.g. axpy) on C++ vectors to accelerate computations to avoid hidden memory allocations/copies [9]. Due to these optimizations, we found that simulating 10 substrates requires approximately 2.6× more computational time than a single substrate [1]. Because the diffusion and decay coefficients are assumed constant, we could pre-compute the forward sweep and part of the back substitution steps for the Thomas solvers, allowing further improvement in the computational speed. [1].

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